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- Applicant: Integrated Genetics, Inc.
 Thew York Avenue
 Framingham Massachusetts 01701(US)
- Inventor: Gordon, Katherine 274 Clarendon Street 14 Boston Massachusetts 02114(US) Inventor: Groet, Suzanne 71 Concord Road Sudbury Massachusetts 01776/US)
- Representative: Deans, Michael John Percy et al Lloyd Wise, Tregear & CO. Norman House 105-109 Strand London WCZR OAE(GB)
- Transgenic animals secreting desired proteins into milk.
- ② A DNA sequence contains a game encoding a protein, the gene being under the transcriptional control in the DNA sequence of a mammalian milk protein promoter which does not naturally control the transcription of the gene, such DNA sequence including DNA enabling secretion of the protein.

FIG I

Xerox Cropy Gentre

EP 0 264 166 A1

TRANSGENIC ANIMALS SECRETING DESIRED PROTEINS INTO MILK

This invention relates to transpenic animals.

It is possible to insert foreign genes into vertebrate embryos, and for these genes to be incorporated into the genome of the resulting animal. Insertion of the foreign genes has been carried out mechanically (by microinjection), and with the aid of retrovirus vectors (for example, as is described in huszar et al. 5 (885) P.N.A.S. U.S.A. 82, 8877). The animals resulting from this process are termed "trensgenic." The foreign genes can be sexually trensmitted through subsequent generations and are frequently expressed in the saimal. In some Instances the proteins encoded by the foreign genes are expressed in specific disease. For example, the metallothineein promoter has been used to direct the expression of the rail growth hormone gene in the liver lissue of transgenic mibe (Palmitter et al., 1982 Nature 303-911), Another example is the electace promoter, which has been shown to direct the expression of foreign genes in the pancrease (Ornitz et al., 1985 Nature 303-901). Developmental control of gene expression has also been activeved in transgenic animals, i.e., the foreign gene is transcribed only during a certain time period, and only in a particular stasse. For example, Magram et al. (1985, Nature 1985,393) demonstrated developmental control of gene under the direction of a glotin promoter; and Krumbauf et al. (1985, Mol. Cell. Biol. §1639) demonstrated developmental control of gene under the direction of a glotin promoter; and Krumbauf et al. (1985, Mol. Cell. Biol. §1639) demonstrated developmental control of gene under the direction of a glotin promoter; and Krumbauf et al. (1985, Mol. Cell. Biol. §1639) demonstrated developmental control of gene expression has also been active and the manufacture of the promoter and the minimum promoters and the promoter and the promoter and the promoter and the direction of a glotin promoter; and Krumbauf et al. (1985, Mol. Cell. Biol. §1639) demonstrated developmental control of gene and the demonstrated developmental control of gene and the promoter and the promoter a

In a first aspect thereof, the invention features a DNA sequence containing a gene encoding a protein, the gene being under the transcriptional control of a mammalian milk protein promoter which does not naturally control the transcription of the gene, the DNA sequence further including DNA enabling secretion in the protein capacity of the protein; a.g., a secretion signal-encoding sequence interposed between the gene and promoter. The promoter can be that of a milk serum protein or a case in protein, although milk serum proteins are preferred, as will be discussed in more detail below. (As used herein, "gene" refers to both genomic DNA sequences and CDNA sequences.)

The invention permits the production of any desired protein in an easily maintained stable, portable culture system, i.e., a living domesticated memmal, which is capable not only of producing the desired as protein, but prelarably of passing on the ability to do so to its female offspring as well. Secretion of the protein into the host mammal's milk facilitates purification and obviates removal of blood products and culture media additives, some of which can be toxic or carcinogenic. More importantly, protein yields will be high and production will be more cost effective and efficient.

The invention thus extends to a mammalian embryo having a nucleus containing a DNA sequence as according to this invention.

According to a second and alternative aspect of this invention, we provide a mammal in which the genome of the mammary glands of said mammal comprises a gene encoding a protein, said gene being under the transcriptional control of a mammalian milk protein promoter which does not naturally control the transcription of said gene, said genome comprising DNA enabling the secretion of said gene encoding said protein.

The invention provides, in a third and further alternative aspect thereof, a method for producing a protein comprising the steps of:

(a) inserting into a mammalian embryo a DNA sequence comprising a gene encoding said protein, said gene being under the transcriptional control of a milk protein promoter which dose not naturally control 40 the transcription of said gene, said DNA sequence comprising DNA enabling secretion of said protein.

(b) allowing said embryo to develop into an adult mammai.

(c) Inducing lactation in said mammal, or in a female descendant of said mammal in which said gene, promoter, and signal sequence are present in the mammary tissue genome.

(d) collecting milk of said lactating mammal, and

(e) isolating said protein from said collected milk.

The invention is hereinafter more particularly described by way of example only with reference to the accompanying drawings, in which:

Fig. I is a diagrammatic representation of the construction of an embodiment of intermediate vector in accordance with the present invention, namely pt-PA VPI-LP(K);

 Fig. 2 is a diagrammatic representation of the construction of another embodiment of intermediate vector in accordance with the invention, namely pWAP (Hs);

Fig. 3 is a diagrammatic representation of the construction of another vector in accordance with the invention, namely pWAP-t-PA(S):

Fig. 4 is a diagrammatic representation of the construction of another intermediate vector in accordance with the invention, namely phibsSVA; and

Fig. 5 is a diagrammatic representation of the construction of another vector according to the invention, namely pWAP-Hbs(S).

s DNA Sequence Elements

Promoter

The milk protein promoter can be derived from any mammalian species, and can be any promoter on naturally associated with any protein which is normally scroted into mammalian milk. Generally, milk proteins which are present in milk in the form of micelles, and which are removed from skim milk by clotting with rennet; and the milk sorm proteins, which are defined herein as the non-caselin milk proteins. Whop proteins constitute the sorum proteins, which are defined herein as the non-caselin milk proteins. Whop proteins constitute the protein wheye acid protein ("WAP") is named based on it is addic isolectic point (Piletz (1991) J Elici. Cham. 256; 18509). Another example of a milk serum protein described in the literature is a-factalbumin (described, along with mouse WAP, in Hennighausen and Sippel (1982 Eur. J. Blocham L25, 13), Milk proteins are discussed in detail in Waistra and Jenness Dairy Chemistry and Physics (John Wiley & Sons 1884).

20 Generally, milk serum protein promoters are preferable to casein promoters in the practice of the present invention

because caseins genorally are produced in female mammals during pregnancy as well as after birth, while WAP is expressed primarily during post-partum lactation. This difference is of potential importance for two reasons. First, pre-birth production of the desired protein under the transcriptional control of a classification of the matter post-parturn. Become, where the desired protein is tonce in large amounts (human tissue plasminogen activator (I-PA) is an exemple), a build-up of the protein in the dissues prior to lactation could be deleterious to line health of the host mammal. An additional advantage of some whey promoters who are the WAP green and the promoter is that they are strong promoters, as evidenced by the large amounts of some whey proteins present in milk. Casein promoters sits have this advantage.

Milk protein genes from which promoters, in addition to the WAP promoters, can be isolated, can be obtained in the same manner in which the WAP genes were isolated, as described in Henrighausen and Sippel, Id. and Campbell et al. (1984) Nucleic Acids Research I2, 8885. The method generally involves isolating by mRNA from a lactating mammary gland, constructing a cDNA library from the mRNA screening like library for the particular milk protein aDNA being sought, cloning that cDNA into vectors, and using the appropriate cDNA as a probe to isolate this genomic clone from a genomic library. A sequence upersean from the irrascription start site in the genomic clone constitutes a puttile "promoter" a genomic sequence proceeding the gener clientest and presumed to be involved in its regulation. The promoter may be isolated by carrying out restriction endourclease dispetions and subloting steps. Promoters and of the of any parfoulder length nor to have directly shown any properties of regulation. The mouse WAP promoters was feetable as a 2.8 to EccRI+ Knyfi fragment immediately 8' to the WAP signal sequence.

Desired Protein

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Any desired protein can be produced according to the invention. Preferred proteins are proteins useful in the treatment, prevention, endoor diagnosis of human disease; examples are LPA and hepatitis B surface antigen. The invention is particularly useful for proteins which must be produced on a large scale to be economical, e.g., industrial enzymes and arisinal proteins.

Signal Sequence

It is necessary, for secretion of the desired protein into the milk of the host mammal, that the DNA sequence containing the pene for the desired protein include DNA which, when transisted, causes the protein to be secreted out of the mammary tissue into the milk. Without such a sequence, the desired protein would remain in the mammary tissue into which purification would be difficult, and would require activities of the host arimals. This DNA can encode a hydrophobic secretion signal which is cleaved during

secretion. The signal sequence can be that which is naturally associated with the desired protein, if the protein is normally secretic (e.g., t-PA). Alternatively, the signal encoding sequence can be that of the milk protein providing the promoter, i.e., when the milk protein gene is digested and the promoter isolated, and fragment is selected which includes both the promoter and the signal encoding sequence directly of ownerterem from the promoter. Another alternative is to employ a signal encoding sequence directly on another secreted protein, which is neither the milk protein normally expressed from the promoter nor the desired nortelin.

to Termination Site

Preferably there is located within or downstream from the 3" end of the desired gene a termination site. This site may be provided by sequences in the gene itself, or may need to be added, if the sequence is to be added, a preferred sequence is provided by the polyadenylation sequence of the virus SV40, as will be described in greater detail below.

Methods

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20 Genetic Manipulations

Generally, all DNA manipulations used in the genetic constructions described herein may be carried out using conventional techniques, as described, e.g., in Meniatis et al. Molecular Cloning: A <u>Laboratory Manual</u> (Cold Spring Harbor Laboratory, 1882).

Introduction of DNA into Embryos

Once the genetic constructions have been produced in vectors, e.g., plasmids, the promoter-signal sequence-desired protein-lermination sequence CNA fragment is excised and then introduced into the desired mammalian embryo using, e.g., retroviruses or standard interorigication methods such as are described in Kreemer et al. (1985), Costantini and Jaenisch, eds., Genetic Mamiguation of the Early Mammalian Embryo, Cold Spring Harbor Laboratory (bowine embryo microinjection); Hammer et al. (1985), Nature 315, 680 (rabbit, sheep, and porcine embryo microinjection); and Gordon and Ruddie (1984) Methods in Embryology (2), 41 (mouse embryo microinjection). Micro-injection is preferably carried out on an embryo at the one-cell stage, to maximize both the chances that the sinjected DNA will be incorporated into all cells of the shimal, including mammary issue, and that the bNA will also be incorporated into the germ cells, so that the similar's offsching with the transport cas well. Microinjection is a standard technique which involves, briefly, isolating fertilized ova, visualizing the pronucleus, and then injecting the DNA into the pronucleus by tholding the ova with a blunt holding pipette of a diameter on the order of 50 Lm, and using a charply pointed pipette of a diameter on the order of 15 Lm to inject buffer-containing DNA into the pronucleus. Following microinjection, the transgenic female animals are allowed to become sexually mature, metad, and milk collected post-parture.

Preferred host mammals are those which are already bred for large volume milk production, e.g., cows. sheep, goats, and pigs

t-PA Production

There will now be described the construction of plasmid DNA in which the gene encoding human uterine LPA, including the signal encoding sequence, is under the transcriptional control of the mouse WAP promoter, and has all its '3' end the SV40 polyadenylation site. This DNA was made from two intermediate plasmids, one carrying the mouse WAP promoter and one carrying the LPA signal and structural sequences, as well as the SV40 polyadenylation site. The WAP promoter containing plasmid pWAP-CAT (Fig. 2, obtained from Lothar Henrighausen. National Institutes of Health) was derived from a plasmid made according to the methods described in Henrighausen and Sippel (BegS; Eur. J. Biotohem. 125, 131 and Campbell et al. (1864) Nucleic Acids Research 12, 8685. In addition to containing the mouse WAP promoter, pWAP-CAT contains a gare which, for present 5 purposes, la tiretevant: the CAT (chloramphenical acetyltransferase) gene, which does not form a part of the final DNA sequence which it emitorisliectade.

Still referring to Fig. 2, pWAP-CAT was modified to convert the Eco Rt site to a Hindflt site using Klenow and Hind III linkers.

The t-PA-containing plasmid pt-PA-VPI-LP(K) (Fig. I) was derived from pt-PAVPI-LP, containing the trPA gene (including the t-PA signal encoding sequence) and SV40 polyadenylation site, by modifying the unique Nool site at the 5 and of the t-PA gene using Nool endoractesse and Klenow and adding Kon linkers to produce a Kont site.

Referring 10 Fig. 3, the <u>Kont-Barn-HI</u> fragment of bc-PA VPH-LPIG, containing the t-PA gene and SV40 secuences, was solated and ligated to <u>Barn-HI-Kprj</u> treated pWAP(H3) to form gWAP-tPA (5), which was 15 then transformed into a TET-ennsitive derivative of <u>E. coll. strain MCI081. This transformed strain, containing plasmid DNA in which the <u>Hirdill-Barn-HI</u> fragment contains the LPA gene including the LPA signal encoding sequence under the transcriptional control of the WAP promoter and followed by the SV40 polyadenylation site, has been deposited in the American Type Culture Collection on March 13, 1986 and given ATCA cossissin No. 67932.</u>

Production of milk into which t-PA has been secreted is carried out by excising the <u>Hindlili-BamHI</u> tragment from the deposited strain and transferring it by microinjection or other means preferably into the one-cell emproy of a mammal ascording to conventional methods, as described above. Alternatively, though less desirably, the entire plasmid or other restriction fragments can be introduced into the embryos are then nutrured to term in vivo. Animals born from such manipulated embryos are screened for 18 the presence of introduced DNA in the genome, and expression of t-PA in the milk is screened for emorg transgenic, leatating females. The protein from the milk of the adult lactating female animal will be assayed for t-PA by conventional procedures.

se Production of Hepatitis B Surface Antigen

Referring to Figure 5, intermediate vectors pWAP-CAT and pHBsSVA were used to construct pWAP-Hbs(5), containing the gene for hepatitis B surface antigen, under the transcriptional control of the WAP promoter and followed by the SV4b polyadonylation site.

The plasmid sWAP-CAT is described above. Plasmid pHbsSVA was constructed as illustrated in Fig. 4, pCLHsA, containing the SV40 pobyadenylation sequence, was restricted with <u>Eco Rt. Sect.</u> and <u>Bglit. pBSBam</u>, containing the gene for hopetitis B surface arrigen, was cut with <u>EcoRt. Bernitia and Pyul</u>, and the two mixtures tigated to give pHbsSVA, in which the SV40 sequence was positioned at the 3' end of the Hbs gene, on a <u>Barniti-Bglit Imgrement. This frampent was then ligitated (Fig. 5) to <u>Barnitial solicitation</u> phosphatase-treated pWAP-CAT, transformed into <u>E. optil</u> strain MCl06i, and the plasmid pWAP-Hbs(S) isolated.</u>

The BamH-EccRI fragment of WAP-Hbs(S) can be excised and used as described enove to produce inputities 5 eurlace antigen. Alternatively, though lass dealrably, the entire pleamed or other restriction fragments can be introduced into the embryos. Embryoe are then nutrured to term in your. Animals born 45 iron such manipulated embryos are screened for the presence of introduced DNA in the genome, and expression of hepatitis 8 surface artigion in the milk is screened for among transperic, lactating formates, pWAP-Hbs(S) has been deposited in the American Type Culture Collection on March 13, 1996 and given ATCC Accression No. 67334. Applicant's assignee, lategrated Genetics, Inc.

Both pWAP-this(S) and pWAP-th-PA(S) can be used as cassette vectors in which the hepatitis B surface antigen given or the t-PA given can be excised and replaced, using conventional methods, with any desired gene. If desired, the signal encoding sequence from pWAP-t-PA(S) can be left in the vector, and a gene lacking such a sequence inserted downstream of and in frame with it. Alternatively, the signal sequence from pWAP-t-PA(S) or pWAP-this(S) can be removed along with the structural gene and the signal encoding sequence of the substituted gene employed. In addition, the WAP promoter alone can be excised and inserted into another desired excreasion vector.

Purification and Use

The proteins produced by a process according to the invention are purified from the milk into which they have been secreted and used for their known purposes.

Hepatitis B surface antigen is useful in the production of hepatitis B vaccine.

1-PA is useful in the treatment of thrombolytic disease in which fibrin clot lysis is necessary, as described in European Patent Application 8530857.3. That Patent Application also describes general purification techniques which will be useful for milk-secreted proteins.

Stability in Milk

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Table I below shows that, despite the presence in milk of numerous proteases, recombinant t-PA is stable when added to raw post milk and ninoutabed at 20° or 3°TC for 24 hours, with no evidence of lose of 75 activity, as measured using the standard fibrin plate lest (results not shown in Table b) or the emidotybic assay described in Wei et al., Id. Similarry, recombinant hepatitis B surface entigen was found to be stable for at least 24 hours in raw goot milk (data not shown).

Table I Amidolytic assay for TPA

					Incubation Time			Temperature	Units/ml	
25	Goat	milk	alone		***			***	<20,	<20
	Goat	milk	&	TPA		0		*** ***	437,	368
	Goat	milk	å	TPA.	24	hours		20°C	419,	434
20	Goat	milk	â	TPA	24	hours.		37°C	. 467.	507

Other Embodiments

Other embodiments are leasible. For example, other milk serum protein promoters can be used in place of the mouse WAP promoter, and the promoter can be derived from any mammalian species. For example, milk serum protein promoters such as that of p-lactoglobulin can be used, and the rat, rather than mouse. WAP promoter can be used; the rat WAP promoter is described in Campbell et al., <u>id</u>. Although less desirable than milk serum protein produced using processes in eccordance with this invention can be any desired protein of therapeutic or industrial importance.

es Claims

- b. A DNA sequence containing a gene encoding a protein, said gene being under the transcriptional control in said DNA sequence of a mammelian milk protein promoter which does not naturally control the transcription of said gene, said DNA sequence further comprising DNA enabling secretion of said protein.
- The DNA sequence of claim I, wherein said secretion-enabling DNA comprises a secretion signal-encoding sequence interposed between said gene and said promoter.
 - 3. The DNA sequence of claim I wherein said milk protein is a milk serum protein or a casein protein.
 - 4. The DNA sequence of claim 3 wherein said milk serum protein is a whey acid protein.
- 5. The DNA sequence of claim T wherein said signal encoding sequence is the signal encoding sequence naturally associated with said gene encoding said protein.
 - The DNA sequence of claim I wherein said signal encoding sequence is the signal encoding sequence naturally associated with said mammalian milk protein promoter.
 - 7. The DNA sequence of claim I wherein said DNA sequence includes a transcriptional stop sequence.

- 8. The DNA sequence of claim 7 wherein said stop sequence is derived from SV40 virus DNA.
- The DNA sequence of claim 7 wherein said stop sequence is contained in the polyedenylation sequence of SV40.
 - 10. A mammalian embryo having a nucleus containing the DNA sequence of claim I.
- II. The DNA sequence of claim I wherein said protein is human fissue plasminogen activator or hepatilis. B surface antiqen,
- 12. A mammal in which the genome of the mammary glands of said mammal comprises a gene encoding a protein, said gene being under the transcriptional control of a mammalian milk protein promoter which does not naturally control the transcription of said gene, said genome comprising DNA enabling the secretion of said gene encoding said protein.
 - 13. The mammal of claim II, said mammal being a sheep, pig, goat, cow, or other mammais.
 - 14. The mammal of claim II wherein said gene is expressed in greater amounts during factation than during pregnancy.
 - 15. A method for producing a protein comprising the steps of:
 - (a) Inserting Into a mammalian embryo a DNA sequence comprising a gene encoding said protein, said gene being under the transcriptional control of a milk protein promoter which does not naturally control the transcription of said gene, said DNA sequence comprising DNA enabling secretion of said posterio.
 - (b) allowing said embryo to develop into an adult mammel,
 - (c) inducing factation in said mammal, or in a female descendant of said mammal in which said gene, promoter, and signal sequence are present in the mammary tissue genome,
 - (d) collecting milk of said factating mammal, and
 - (e) isolating said protein from said collected milk.

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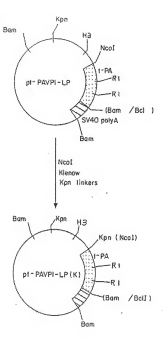


FIG I





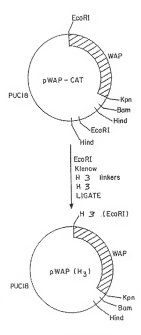


FIG 2



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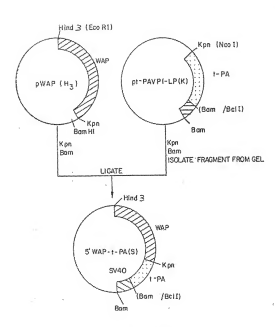
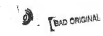


FIG 3



Neu eingessicht (* Lusty Stad) Neuvollument dépasé

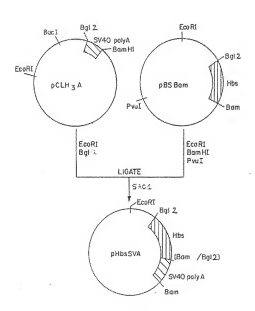


FIG 4

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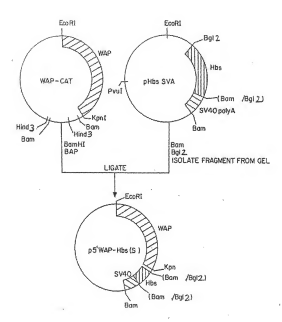


FIG 5



EUROPEAN SEARCH REPORT

Category A	of relea	h indication, where appropriate, ant passages	Relevant	CLASSIFICATION OF THE
Α	EP - A1 - 0 169 6		to claim	APPLICATION (Int. CL4)
	FELLOWS OF HARVAI	572 (PRESIDENT AND	1,10, 12,15	C 12 N 15/00
	* Claims 1,4,1	13 *		C 07 H 21/04
	***	•		C 12 P 21/00
D,A	NATURE, vol. 313, New York, London	10,12		
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	* Totality *			
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D,A	NATURE, vol. 315, New York, London	May 23, 1985,	10,12	
	J. MAGRAM et al. regulation of clo gene in transgens pages 338-340	ned adult B-globin		TECHNICAL FIELDS SEARCHED (IVI. C: 4)
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D,A	NATURE, vol. 300, 16, 1982, New Yor	10,12		
	R.D.PALMITER et s bf mice that deve microinjected wit growth hormone fu pages 611-615	To the state of th		
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***************************************	The present search report has t	where drawers up for all claims		***************************************
***********	Place of exercit	Date of completion of the search	1	Examiner
	VIENNA	25-06-1987		WOLF
Y:px	CATEGORY OF CITED DOCT articularly relevant if taken alone articularly relevant if combined to ocument of the same category ichnological background on-written disclosure	E: sertier pate after the fill O: document L: document	nt document ng date sited in the a sited for othe	riying the invention , but published on, or pplication ir reasons tent lamily, oprresponding

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	DOCUMENTS CONS	DERED TO BE RE	EVANT.		EP 87303112.4	
Category	Citation of document with of refers	indication, where appropris int passages	4,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)	
D,A	MOLECULAR AND CEN	LULAR BIOLOGY	, vol n DC	10,12		
	R. KRUMLAUF et al Regulation of √-1 in Transgenic Mic pages 1639-1648	etoprotein Ge				
	* Totality *					
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D,A	EUROPEAN JOURNAL vol. 125, no. 1, Berlin, Heidelber	June 2, 1982,	RY,	1,3	***************************************	
	L.G.HENNIGHAUSEN terization and Cl Specific for the Mammary Gland" pages 131-141	oning of the	mRNAs			
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	The present search report has b	awn drawn up for all claims				
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	VIENNA	25-06-19	87		WOLF	
Y: pa	CATEGORY OF CITED DOCU recuising relevant if taken alone inscularly relevant if combined we cument of the same category chnological background in-written disclosure termediate document	ith another 0:	T: theory or principle underlying the invention E: earlier pattern document, but published on, or after the filling date and only or L document clied in the application L document clied in the application L document clied for other reasons ā: member of the same patent family, corresponding document			